



INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT) WO 99/00113 (11) International Publication Number: (51) International Patent Classification 6: $\mathbf{A1}$ 7 January 1999 (07.01.99) A61K 9/22 (43) International Publication Date: David, C. [US/US]; 13626 Franklin Street #3, Whittier, CA PCT/US98/13272 (21) International Application Number: 90602 (US). 26 June 1998 (26.06.98) (74) Agent: RAYMER, Gregory, P.; Gray Cary Ware & Freidenrich, (22) International Filing Date: Suite 1600, 4365 Executive Drive, San Diego, CA 92121 (30) Priority Data: US 27 June 1997 (27.06.97) 60/051,021 9 September 1997 (09.09.97) US (81) Designated States: AL, AM, AT, AU, AZ, BA, BB, BG, BR, 08/926,155 BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GE, GH, GM, GW, HU, ID, IL, IS, JP, KE, KG, KP, KR, KZ, (63) Related by Continuation (CON) or Continuation-in-Part

60/051,021 (CIP)

08/926,155 (CIP)

27 June 1997 (27.06.97)

9 September 1997 (09.09.97)

(71) Applicant (for all designated States except US): VIVORX PHARMACEUTICALS, INC. [US/US]; 2825 Santa Monica Boulevard, Santa Monica, CA 90404 (US).

(CIP) to Earlier Applications

(72) Inventors; and

US

US

Filed on

Filed on

(75) Inventors/Applicants (for US only): DESAI, Neil, P. [IN/US]; 3633 Purdue Avenue, Los Angeles, CA 90066 (US). SOON-SHIONG, Patrick [US/US]; 11755 Chenault Street, Los Angeles, CA 90049 (US). MAGDASSI, Shlomo [IL/IL]; Hanerd Street 36, Jerusalem (IL). SAHADEVAN,

LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZW, ARIPO patent (GH, GM, KE, LS, MW, SD, SZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European

patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG).

Published

With international search report.

(54) Title: NOVEL FORMULATIONS OF PHARMACOLOGICAL AGENTS, METHODS FOR THE PREPARATION THEREOF AND METHODS FOR THE USE THEREOF

(57) Abstract

In accordance with the present invention, there are provided compositions and methods useful for the in vivo delivery of substantially water insoluble pharmacologically active agents (such as the anticancer drug paclitaxel) in which the pharmacologically active agent is delivered in the form of suspended particles coated with protein (which acts as a stabilizing agent). In particular, protein and pharmacologically active agent in a biocompatible dispersing medium are subjected to high shear, in the absence of any conventional surfactants, and also in the absence of any polymeric core material for the particles. The procedure yields particles with a diameter of less than about 1 micron. The use of specific composition and preparation conditions (e.g., addition of a polar solvent to the organic phase), and careful selection of the proper organic phase and phase fraction, enables the reproducible production of unusually small nanoparticles of less than 200 nm diameter, which can be sterile-filtered. The particulate system produced according to the invention can be converted into a redispersible dry powder comprising nanoparticles of water-insoluble drug coated with a protein, and free protein to which molecules of the pharmacological agent are bound. This results in a unique delivery system, in which part of the pharmacologically active agent is readily bioavailable (in the form of molecules bound to the protein), and part of the agent is present within particles without any polymeric matrix therein.